

**IN THE UNITED STATES PATENT AND TRADEMARK OFFICE**

In re application of: Wachtmann

Application No.: 10/670,673

Filed: September 25, 2003

For: METHOD OF FORMING A SURFACE MICROMACHINED MEMS DEVICE

Group No.: 2814

Examiner: Pizarro Crespo, Marcos D

**Mail Stop Appeal Briefs – Patents**

**Commissioner for Patents**

**P.O. Box 1450**

**Alexandria, VA 22313-1450**

**TRANSMITTAL OF SUPPLEMENTAL REPLY BRIEF**

1. Transmitted herewith is the SUPPLEMENTAL REPLY BRIEF in this application with respect to the Examiner's Supplemental Answer mailed on May 12, 2008.

**STATUS OF APPLICANT**

2. This application is on behalf of a large entity.

**EXTENSION OF TERM**

3. Applicant believes that no extension of term is required. However, this conditional petition is being made to provide for the possibility that applicant has inadvertently overlooked the need for a petition and fee for extension of time.

Charge any fees required by this paper to deposit account number 19-4972.

Date: June 3, 2008

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant:	Wachtmann	Att. Docket:	2550/185
Serial No.:	10/670,673	Art Unit:	2814
Filing Date:	September 25, 2003	Examiner:	Pizarro Crespo, Marcos D
Invention:	METHOD OF FORMING A SURFACE MICROMACHINED MEMS DEVICE		

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**SUPPLEMENTAL REPLY BRIEF**

Applicant submits this supplemental reply brief for the subject application, in response to the examiner's supplemental answer ("ESA"), which was mailed on May 12, 2008. This ESA was in response to the remand by the Board of April 17, 2008, for further consideration of a rejection.

In Applicant's Reply Brief, Applicant stated in pertinent part:

"If Montague's silicon nitride insulating layer 22 is replaced by an oxide layer, as taught by Kim or Lee, the etchant material used to dissolve the sacrificial layers 30, 32 will also dissolve the oxide in insulating layer 22. Dissolution of insulating layer 22 yields a non-functional device because this layer 22 will de-bond from the substrate allowing the conductive layer 24 to short with the substrate. Thus, claim 1 is non-obvious in view of Montague/Kim/Lee because the process taught by the combination of references will not form a properly functioning MEMS device. "

In the ESA, the Examiner responded in pertinent part:

"..., Kim clearly teaches that aluminum oxide has an etching selectivity of about 5 with respect to silicon dioxide. In other words, the sacrificial layers of Montague would etch five times faster than the oxide layer of Kim and Montague. See, e.g., col.3/11.25-26 and col.4/11.63-66. Thus, the etching material used to dissolve the sacrificial layers 30, 32 of Montague will not dissolve the oxide material suggested by Kim and Lee, as the appellants are arguing above."

Applicant respectfully traverses the Examiner's conclusion, as derived by the Examiner from the disclosure of Kim '763, for the following reasons:

1. Montague '283's process uses a **wet etch** to dissolve sacrificial layers 30 and 32 to release the MEMS structures. (See Montague '283, col. 8, lines 59-61). These sacrificial layers are formed of silicon dioxide or silicate glasses. (See Montague '283, col. 5, lines 64-66).

2. Kim '763's teachings about aluminum oxide as an etch stop and statements about etching selectivity relative to silicon oxide are for a **dry etch** process only. (See, Kim '763, col. 4, lines 63-66.) The Examiner's assertion that "... the sacrificial layers of Montague would etch five times faster than the oxide layer of Kim and Montague" simply does not apply to Montague's wet etch process.

3. For a **wet etch** process, using a typical wet etchant, such as 5:1 BHF, evaporated aluminum oxide dissolves at a faster rate than Montague's silicon dioxide sacrificial layers 30 and 32. (In 5:1 BHF, evaporated aluminum oxide etches at 160 nm/min. versus thermal silicon dioxide, which etches at 100 nm/min.)

As stated in Applicant's Reply Brief, the wet etchant material used to dissolve Montague's sacrificial layers 30, 32 will, therefore, also dissolve aluminum oxide if used in insulating layer 20. Dissolution of insulating layer 20 yields a non-functional device because this layer 20 will de-bond from the substrate allowing the conductive layer 24 to short with the substrate. In contrast, silicon nitride, deposited by the low pressure chemical vapor deposition technique preferred by Montague, has a slow or zero etch rate when exposed to wet etchant

5:1 BHF. Therefore, silicon nitride is suitable both as an etch stop and as an insulating layer, while aluminum oxide is not. Clearly, no one skilled in the art would consider substituting aluminum oxide for silicon nitride as the insulating layer 20 in Montague's process.

Please note that the etch rates cited in the preceding paragraph are well known to those skilled in the art. For example, these etch rates are given in tables III and IV from "Etch Rates for Micromachining Process, Part II," Kirt R. Williams and Kishan Gupta, *IEEE Journal of Microelectromechanical Systems*, vol. 12, no. 6, (December 2003). A copy of this paper is attached for reference.

For the reasons set forth in Applicant's Appeal Brief, Reply Brief and those expressed above, Applicant submits that all pending claims in the application are allowable over the art of record and early notice to that effect is respectfully solicited.

Respectfully submitted,

Date: June 3, 2008

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# Etch Rates for Micromachining Processing—Part II

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**Abstract**—Samples of 53 materials that are used or potentially can be used in the fabrication of microelectromechanical systems and integrated circuits were prepared: single-crystal silicon with two doping levels, polycrystalline silicon with two doping levels, polycrystalline germanium, polycrystalline SiGe, graphite, fused quartz, Pyrex 7740, nine other preparations of silicon dioxide, four preparations of silicon nitride, sapphire, two preparations of aluminum oxide, aluminum, Al<sub>2</sub>/Si, titanium, vanadium, niobium, two preparations of tantalum, two preparations of chromium, Cr on Au, molybdenum, tungsten, nickel, palladium, platinum, copper, silver, gold, 10 Ti/90 W, 80 Ni/20 Cr, TiN, four types of photoresist, resist pen, Parylene-C, and spin-on polyimide. Selected samples were etched in 35 different etches: isotropic silicon etchant, potassium hydroxide, 10:1 HF, 5:1 BHF, Pad Etch 4, hot phosphoric acid, Aluminum Etchant Type A, titanium wet etchant, CR-7 chromium etchant, CR-14 chromium etchant, molybdenum etchant, warm hydrogen peroxide, Copper Etchant Type CE-200, Copper Etchant APS 100, dilute aqua regia, AU-5 gold etchant, Nichrome Etchant TFN, hot sulfuric+phosphoric acids, Piranha, Microstrip 2001, acetone, methanol, isopropanol, xenon difluoride, HF+H<sub>2</sub>O vapor, oxygen plasma, two deep reactive ion etch recipes with two different types of wafer clamping, SF<sub>6</sub> plasma, SF<sub>6</sub>+O<sub>2</sub> plasma, CF<sub>4</sub> plasma, CF<sub>4</sub>+O<sub>2</sub> plasma, and argon ion milling. The etch rates of 620 combinations of these were measured. The etch rates of thermal oxide in different dilutions of HF and BHF are also reported. Sample preparation and information about the etches is given. [1070]

**Index Terms**—Chemical vapor deposition (CVD), etching, evaporation, fabrication, materials processing, micromachining.

## I. INTRODUCTION

WHEN designing a microfabrication process, the etch rate of each material to be etched must be known. Knowing the etch rates of other materials that will be exposed to the etch, such as masking films and underlying layers, enables an etch process to be chosen for good selectivity (high ratio of etch rate of the target material to etch rate of the other material)—if one exists. While several large literature-review compilations of etches that target specific materials have been made [1], [2], these only report etch rates in some cases, and rarely have corresponding selectivity information. This paper provides such information, expanding on an earlier paper [3] to give 620 etch rates of 53 materials in 35 etches that have been used or may

be used in future fabrication of microelectromechanical systems (MEMS) and integrated circuits (ICs) (approximately 50 etch rates measured in the earlier paper have been included in this one). These data allow the selection of new combinations of structural material, underlying material, and etchant for micro-machining.

Table I summarizes the etches tested, abbreviated names for the etches, and the target materials for each. Table II lists etch rates of Si, Ge, SiGe, and C in the SI units of nm/min (not Å/min as in the earlier tables) [3]. Table III covers films and wafers that are primarily silicon dioxide, produced under many different conditions. Table IV is on silicon nitride and aluminum oxide. Table V covers the metals Al, Ti, V, Nb, Ta, and Cr. Table VI continues with the metals Mo, W, Ni, Pd, Pt, Cu, Ag, Au, alloys 10 Ti/90 W, 80 Ni/20 Cr, and compound TiN. Finally, Table VII gives etch rates of organics: photoresists, a resist pen, and a spin-on polyimide.

Section II of this paper lists the materials etched, their preparation, and some uses or potential uses in MEMS and ICs. Section III describes the preparation and applications of the wet and dry etches that were studied, as well as some key experimental results. Section IV describes etch-rate measurement techniques, and Section V discusses the results.

## II. SAMPLE PREPARATION

The preparation of the samples in the etch-rate tables is described below, listed by the labels (in *italics*) used across the tops of the tables. All coated materials were deposited on 100-mm-diameter silicon wafers. For the isotropic silicon etchant, potassium hydroxide, and a few other etches, the wafers were first coated with LPCVD silicon nitride so that etches would not penetrate into the silicon or attack the back side of the wafer.

In several cases, similar materials were prepared using different methods (e.g., wafer form, PECVD, LPCVD, and ion-milled silicon dioxide; annealed and unannealed films) to study and emphasize the effect on their etching characteristics.

Existing or potential MEMS applications are given for the materials. Many of the materials were discussed in more detail previously [3].

### A. Silicon, Germanium, SiGe, and Carbon

*(100) Si Low-Doped Wafer:* Single-crystal silicon, (100) orientation, phosphorus-doped n-type, resistivity of 3–40 Ω-cm, grown with the Czochralski (CZ) method. Single-crystal silicon is the standard starting material for bulk micromachining.

*Float-Zone Si Wafer:* Single-crystal silicon, (100) orientation, undoped, grown with the float-zone (FZ) method for a high resistivity of >10 000 Ω-cm. Float-zone wafers have been used as substrates in RF MEMS application to reduce eddy-current loss.

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TABLE I  
ETCH DESCRIPTIONS, ABBREVIATIONS, AND TARGET MATERIALS

Etchant	Etchant Abbrev.	Target Material
Isotropic Silicon Etchant "Trilogy Etch" (126 HNO <sub>3</sub> : 60 H <sub>2</sub> O : 5 NH <sub>4</sub> F), -20°C	Si Iso Etch	Silicon
KOH (30% by weight), 80°C	KOH	Silicon QDE
10:1 HF (5 H <sub>2</sub> O : 1 49% HF), -20°C	10:1 HF	Silicon Dioxide
5:1 BHF (5 40% NH <sub>4</sub> F : 1 49% HF), -20°C	5:1 BHF	Silicon Dioxide
Pad Etch 4 from Ashland (13% NH <sub>4</sub> F + 32% HAc + 49% H <sub>2</sub> O + 6% propylene glycol + surfactant), -20°C	Pad Etch 4	SiO <sub>2</sub> , not Al
Phosphoric Acid (85% by weight), 160°C	Phosphoric	Silicon nitride
Al Etchant Type A from Transene (80% H <sub>3</sub> PO <sub>4</sub> + 5% HNO <sub>3</sub> + 5% HAc + 10% H <sub>2</sub> O), 50°C	Al Etch A	Aluminum
Titanium wet etchant (20 H <sub>2</sub> O : 1 H <sub>2</sub> O <sub>2</sub> : 1 HF), -20°C	Ti Etch	Titanium
Chromium etchant CR-7 from Cyanatek (9% (NH <sub>4</sub> ) <sub>2</sub> Ce(NO <sub>3</sub> ) <sub>6</sub> + 6% HClO <sub>4</sub> + H <sub>2</sub> O), -20°C	CR-7	Chromium
Chromium etchant CH-14 from Cyanatek (22% (NH <sub>4</sub> ) <sub>2</sub> Ce(NO <sub>3</sub> ) <sub>6</sub> + 8% HAc + H <sub>2</sub> O), -20°C	CR-14	Chromium
Molybdenum etchant (180 H <sub>3</sub> PO <sub>4</sub> : 11 HAc : 11 HNO <sub>3</sub> : 150 H <sub>2</sub> O), -20°C	Moly Etch	Molybdenum
Hydrogen peroxide (30wt% H <sub>2</sub> O <sub>2</sub> : 70wt% H <sub>2</sub> O), 50°C	H <sub>2</sub> O <sub>2</sub> 50°C	Tungsten
Copper etchant type CE-200 from Transene (30% FeCl <sub>3</sub> + 3-4% HCl + H <sub>2</sub> O), -20°C	Cu FeCl <sub>3</sub> 200	Copper
Copper etchant APS 100 from Transene (15-20% (NH <sub>4</sub> ) <sub>2</sub> S <sub>2</sub> O <sub>8</sub> + H <sub>2</sub> O), 30°C	Cu APS 100	Copper
Dilute aqua regia (3 HCl : 1 HNO <sub>3</sub> : 2 H <sub>2</sub> O), -30°C	Dil. Aqua regia	Noble metals
Gold etchant AU-5 from Cyanatek (5% I <sub>2</sub> + 10% KI + 85% H <sub>2</sub> O), -20°C	AU-5	Gold
Nichrome etchant TFN from Transene (10-20% (NH <sub>4</sub> ) <sub>2</sub> Ce(NO <sub>3</sub> ) <sub>6</sub> + 5-6% HNO <sub>3</sub> + H <sub>2</sub> O), -20°C	NiCr TFN	NiCr
1 H <sub>2</sub> SO <sub>4</sub> : 1 H <sub>3</sub> PO <sub>4</sub> : 160°C	Phos+Sulf	Sapphire
Piranha (-50 H <sub>2</sub> SO <sub>4</sub> : 1 H <sub>2</sub> O <sub>2</sub> ), 120°C	Piranha	Cleaning
Microslip 2001 photoresist stripper, 85°C	Microslip	Photoresist
Acetone, -20°C	Acetone	Photoresist
Methanol, -20°C	Methanol	Cleaning
Isopropanol, -20°C	IPA	Cleaning
XeF <sub>2</sub> , 2.6 mTorr, homemade chamber	XeF <sub>2</sub>	Silicon
HF + H <sub>2</sub> O vapor, 1 cm over dish with 49% HF	HF vapor	Silicon dioxide
Technics plasma, O <sub>2</sub> , 400 W @ 30 kHz, 300 mTorr	Technics O <sub>2</sub>	Photoresist
STS ASE DRIE, mechanical chuck, high frequency, typical recipe	DRIE HF mech.	Silicon
STS ASE DRIE, electrostatic chuck, high frequency, typical recipe	DRIE HF ES	Silicon
STS ASE DRIE, mechanical chuck, stop-on-oxide (low-frequency platen), typical recipe	DRIE LF mech	Silicon
STS ASE DRIE, electrostatic chuck, stop-on-oxide (low-frequency platen), typical recipe	DRIE LF ES	Silicon
STS 320 RIE, SF <sub>6</sub> , 100 W @ 13.56 MHz, 20 mTorr	STS 320 SF <sub>6</sub>	Si, SiN, metals
STS 320 RIE, SF <sub>6</sub> + O <sub>2</sub> , 100 W @ 13.56 MHz, 20 mTorr	STS SF <sub>6</sub> +O <sub>2</sub>	Si, SiN, metals
STS 320 RIE, CF <sub>4</sub> , 100 W @ 13.56 MHz, 60 mTorr	STS 320 CF <sub>4</sub>	Si, SiO <sub>2</sub> , SiN
STS 320 RIE, CF <sub>4</sub> + O <sub>2</sub> , 100 W @ 13.56 MHz, 60 mTorr	STS CF <sub>4</sub> +O <sub>2</sub>	Si, SiO <sub>2</sub> , SiN
Ion milling with argon ions at 500 V, -1 mA/cm <sup>2</sup> , normal incidence (Commonwealth data)	Ion Mill	Everything

Notation:

BHF = buffered hydrofluoric acid

DRIE = deep reactive ion etch

RIE = reactive ion etch

STS ASE = Surface Technology Systems Advanced Silicon Etch

**Polysilicon LPCVD Undoped:** Undoped polycrystalline silicon deposited in a Tylan low-pressure chemical-vapor deposition (LPCVD) furnace with recipe SiH<sub>4</sub> flow rate = 100 sccm, temperature (T) = 605 °C, pressure (p) = 300 mTorr. Deposited on a wafer with 100 nm of thermal oxide on it to enable interferometric thickness measurements. Refractive index (RI) = 3.97. Undoped poly, which has a high sheet resistance as deposited, is the most common structural material for surface micromachining. It can be doped with ion implantation or by diffusing in dopant atoms from an adjacent film (e.g., PSG, below) at high temperature.

**Polysilicon LPCVD In-Situ n<sup>+</sup>:** An n-type, phosphorus-doped polycrystalline silicon deposited in a Tylan LPCVD furnace with recipe SiH<sub>4</sub> = 100 sccm, 1.6% PH<sub>3</sub>/balance SiH<sub>4</sub> = 2 sccm, T = 610 °C, p = 300 mTorr. Deposited on a wafer with thermal oxide on it to enable interferometric thickness measurements. RI = 3.97. *In situ* doping gives a conducting film, useful for thicker films and in cases in which other considerations limit the temperature. The deposition rate is about 1/3 that of undoped polysilicon under similar conditions.

**Poly Ge LPCVD Undoped:** Undoped polycrystalline germanium deposited in Tystar LPCVD furnace with recipe GeH<sub>4</sub> =

TABLE II  
 ETCH RATES OF Si, Ge, SiGe, AND C (nm/min)

Etch	Si (100) Water	Floater Zone Si Water	Poly Si LPCVD Undoped	Poly Si LPCVD In-situ n <sup>+</sup>	Poly Ge LPCVD Undoped	Poly SiGe LPCVD P-type	Graphite Ion-Milled
Si Iso Etch	150	W	100	310	890	550	60
KOH	1100	F	870	>1000	-	-	-
10:1 HF	S	S	0	0.7	0	0.42	-
5:1 BH-F	0	S	0.2	0.9	R 1.8	0.45	R 17
Pad Etch 4	S	S	S	S	-	-	-
Phosphoric	0.17	S	S	0.7	0.13	0.40	-
Al Etch A	S	S	<0.9	<1	13	0.11	-
Ti Etch	S	S	S	1.2	-	-	-
CR-7	0	S	0	S	260	0.35	< 0.5
CR-14	S	S	0	S	-	-	-
Moly Etch	-	-	-	-	-	-	-
H <sub>2</sub> O <sub>2</sub> 50°C	S	S	S	S	460	0.13	-
Cu FeCl <sub>2</sub> 200	-	-	-	-	-	-	-
Cu APS 100	-	-	-	-	-	-	-
Dil. Aqua regia	0	0	0	0	-	-	-
AJ-5	S	S	0	S	-	-	-
NiCr TFN	0	S	S	S	-	-	-
Phos+Sn	0.88	S	S	S	-	-	-
Piranha	0	S	0	0	soft	0	-
Microship	S	S	S	S	-	-	0
Acetone	S	S	0	0	-	-	-
Methanol	S	S	0	S	-	-	S
IPA	S	S	S	S	-	-	S
XeF <sub>2</sub>	460	W	160	190	-	-	-
HF vapor	S	S	0	0	-	-	-
Technics O <sub>2</sub>	S	S	0	0	-	-	0
DRIE HF mach.	1500	1600	W	W	-	-	-
DRIE HF ES	2400	W	W	W	400	1400	-
DRIE LF mach.	2400	W	W	W	-	-	-
DRIE LF ES	2000	W	W	W	170	1040	-
STS 320 SF <sub>6</sub>	W	W	W	W	-	-	-
STS SF <sub>6</sub> +O <sub>2</sub>	1500	W	W	W	-	-	-
STS 320 CF <sub>4</sub>	W	W	W	W	-	-	-
STS CF <sub>4</sub> +O <sub>2</sub>	95	-	-	-	-	-	-
Ion Mill	36	36	36	36	-	-	4.4

Notation:

W = Etch known to work, but etch rate not measured

F = Etch known to be fast, but etch rate not measured

S = Etch rate known to be slow or zero, but etch rate not measured

R = Film was visibly roughened or attacked

T = Thicker after etch (due to swelling or compound formation)

P = Some of film peeled during etch or when rinsed

I = Incubation time before etching fully starts

C = Film coagulated

soft = a soft material remained after etching

Ion-Milled = Ion-mill deposited

LPCVD = low-pressure chemical-vapor deposition

poly Ge = polycrystalline germanium

poly SiGe = polycrystalline SiGe

poly Si = polycrystalline silicon

100 sccm,  $T = 400^\circ\text{C}$ ,  $p = 300$  mtorr. The polygermanium deposition was preceded by the deposition of silicon seed layer approximately 6 nm thick using the recipe  $\text{Si}_2\text{H}_6 = 100$  sccm,  $T = 400^\circ\text{C}$ ,  $p = 300$  mtorr.

Germanium forms an oxide that is soluble in water. Thus, water with a high concentration of dissolved oxygen etches germanium. Hydrogen peroxide is a useful etchant for Ge, etching faster at higher temperature.

Polygermanium has been used in surface micromachining as a sacrificial layer in conjunction with a polycrystalline SiGe structural layer, using warm hydrogen peroxide as the etchant [4]. The relatively low deposition temperatures are compatible with CMOS circuitry with aluminum interconnections.

**Poly SiGe LPCVD p-type:** A p-type polycrystalline silicon-germanium deposited in a Tysar LPCVD furnace with recipe  $\text{SiH}_4 = 100$  sccm,  $\text{GeH}_4 = 58$  sccm,  $\text{B}_2\text{H}_6 = 60$  sccm,  $T =$



TABLE III  
ETCH RATES OF SILICON DIOXIDE (nm/min)

Etch	Fused Quartz	Pyrex 7740	Thermal Oxide	Ann LTO	Unan LTO	Ann LTO	Unan LTO	Ann PSG	Oxide	Oxide	Oxide
	Water	Water	Wet-Gm	Calogic	Tylan	Tylan	Tylan	Tylan	PECVD	PECVD	Ion-Mill
Si Iso Etch	12	R 140	8.7	15	-	11	400	170	100	25	43
KOH	8.7	11	7.7	8.1	-	9.4	-	38	15	7.8	6.0
10:1 HF	26	W	23	W	W	34	1500	470	W	W	W
5:1 BHF	130	43	100	150	W	120	690	440	450	240	82
Pad Etch 4	19	17	31	W	38	W	230	W	160	W	W
Phosphoric	0.23	3.7	0.18	S	0.21	2.7	1.8	-	-	-	S
Al Etch A	S	-	0	S	S	0	S	< 1	0	S	S
Ti Etch	-	-	12	W	W	W	W	210	W	W	-
CF-7	R < 0.4	R 0	0.02	0	S	S	S	S	0	0	0
CR-14	S	-	0.01	S	S	S	S	S	S	S	S
Moly Etch	-	-	-	-	-	-	-	-	-	-	-
H <sub>2</sub> O <sub>2</sub> 52°C	S	S	0	S	S	S	S	S	S	S	S
Cu FeO <sub>2</sub> 200	-	-	-	-	-	-	-	-	-	-	-
Cu APS 100	-	-	-	-	-	-	-	-	-	-	-
DI Aqua regia	0	-	0	0	0	0	0	0	0.7	S	S
AU-5	S	-	S	S	S	S	S	S	0	S	S
WCl <sub>6</sub> TRN	S	-	S	S	S	S	S	S	S	S	S
Phos-Sulf	S	-	0.057	S	S	S	S	S	S	S	S
Piranha	R 0	R 0	0	0	S	0	S	0	0	0	0
Microstrip	S	S	S	S	S	S	S	S	S	S	S
Acetone	S	S	0	S	S	0	S	0	S	S	S
Methanol	S	S	S	S	S	S	S	S	S	S	S
IPA	S	S	S	S	S	S	S	S	S	S	S
XeF <sub>4</sub>	S	-	0	S	S	0	0	0	0	S	S
HF vapor	W	W	66	W	W	78	210	150	W	W	W
Technics O <sub>2</sub>	S	S	0	S	S	0	0	0	S	S	S
DRIE HF mech.	S	-	S	7.5	8.2	6.9	9.5	11	9.5	S	S
DRIE HF ES	S	-	24	W	W	W	W	W	W	W	W
DRIE LF mech.	S	-	S	3.6	9.8	9.4	15	15	4.0	S	S
DRIE LF ES	S	-	24	W	W	W	W	W	W	W	W
STS 320 SF <sub>6</sub>	W	-	W	W	W	W	W	W	W	W	W
STS SF <sub>6</sub> +O <sub>2</sub>	30	10	29	38	58	48	73	60	55	32	30
STS 320 CF <sub>4</sub>	W	W	W	30	W	W	W	W	W	W	W
STS CF <sub>4</sub> +O <sub>2</sub>	41	31	44	42	51	46	69	62	51	43	21
Ion Mill	W	W	39	W	W	W	W	W	W	W	W

Notation:

W = Etch known to work, but etch rate not measured  
 F = Etch known to be fast, but etch rate not measured  
 S = Etch rate known to be slow or zero, but etch rate not measured  
 R = Film was visibly roughened or attacked  
 T = Thicker after etch (due to swelling or compound formation)  
 P = Some of film peeled during etch or when rinsed  
 I = Incubation time before etching fully starts  
 C = Film coagulated  
 soft = a soft material remained after etching

Ann LTO = annealed low-temperature oxide  
 Ann. PSG = annealed phosphosilicate glass  
 LPCVD = low-pressure chemical vapor deposition  
 PECVD = plasma-enhanced chemical-vapor deposition  
 Unan LTO = unannealed low-temperature oxide  
 Unan. PSG = unannealed phosphosilicate glass

100 °C,  $p = 300$  mtorr. This film is approximately 48 atomic % Ge.

**Graphite Ion-Milled:** Graphite ion-mill-deposited (also known as ion-beam-deposited) in a Commonwealth Scientific system from a graphite target with argon ions at 1250 V, current density of about 2 mA/cm<sup>2</sup> (~200 mA over most of a 5-inch-diameter target), chamber  $p = 3 \times 10^{-4}$  torr. Graphite has had little or no use in MEMS to date. In this work, it was found to be easily deposited and etched in silicon isotropic etchant. It may find use as a hard mask for plasma etching due to its low etch rate, and as a dry lubricant in MEMS.

## B. Silicon Dioxide

**Fused Quartz Wafer:** Wafers of General Electric 124 or NSG N fused quartz source material, >99.99% silicon dioxide, with amorphous structure (as opposed to true crystalline quartz). This material is commonly referred to simply as "quartz." It is compatible with silicon-wafer processing steps, and may find application as a substrate in RF MEMS as it is not conductive, eliminating eddy-current losses.

**Pyrex 7740 Wafer:** Corning Pyrex 7740 glass, 81% SiO<sub>2</sub>, 13% B<sub>2</sub>O<sub>3</sub>, 4% Na<sub>2</sub>O, 2% Al<sub>2</sub>O<sub>3</sub>. Pyrex 7740 (and the very

TABLE IV  
 ETCH RATES OF SILICON NITRIDE AND ALUMINUM OXIDE (nm/min)

Etch	Stoich Si Nit. LPCVD	Si-Rich Si Nit. LPCVD	PECVD Silicon Nit. Low RI	PECVD Silicon Nit. High RI	Sapphire Water	Aluminum Oxide Ion-Mill	Aluminum Oxide Evap
Si Iso Etch	-	0.23	> 66	12	R < 0.7	99	12
KOH	0	0	0.67	0	R 0	> 2500	> 800
10:1 HF	1.1	S	-	-	S	-	-
5:1 BHF	S	1.3	60	8.2	0	-	160
Pad Etch 4	0.41	S	-	1.6	S	-	-
Phosphoric	4.5	2.7	W	20	< 0.1	-	> 5
Al Etch A	S	< 0.05	-	-	R < 2	65	5.7
Tr Etch	0.99	S	-	-	S	-	-
Cr-7	S	0	< 0.14	0	R 0	0.34	0.075
Cr-14	S	S	-	-	S	-	-
Moly Etch	-	-	-	-	-	-	-
H <sub>2</sub> O <sub>2</sub> 50°C	0	S	S	S	S	-	-
Cu FeCl <sub>3</sub> 200	-	0	-	-	-	-	-
Cu APS 100	-	0	-	-	-	-	-
Dil. Aqua regia	0	0	-	-	S	-	1.1
AU-5	S	0	-	-	-	-	-
NiCr TFN	S	S	-	-	-	-	-
Phos+Sulf	2.9	S	W	10	< 0.3	-	-
Piranha	0	0	< 0.04	0	R 0	97	19
Microstrip	S	S	S	S	S	-	-
Acetone	0	0	S	S	S	0	S
Methanol	S	0	S	S	S	0	S
IPA	S	0	S	S	S	0	S
XeF <sub>2</sub>	12	-	-	-	-	-	-
HF vapor	1.0	1.9	-	-	S	-	-
Technics O <sub>2</sub>	0	S	S	S	S	S	S
DRIE HF mech.	W	21	W	W	S	S	S
DRIE HF ES	W	W	W	W	S	S	S
DRIE LF mech.	W	26	W	W	S	S	S
DRIE LF ES	W	W	W	W	S	S	S
STS 320 SF <sub>5</sub>	W	W	W	W	S	S	S
STS SF <sub>4</sub> +O <sub>2</sub>	150	150	200	190	2.2	0.55	0.41
STS 320 CF <sub>4</sub>	34	W	W	W	S	S	S
STS CF <sub>4</sub> +O <sub>2</sub>	120	> 130	240	110	0	< 2	< 0.2
Ion Mill	13	9.4	W	W	W	10	10

Notation:

W = Etch known to work, but etch rate not measured

F = Etch known to be fast, but etch rate not measured

S = Etch rate known to be slow or zero, but etch rate not measured

R = Film was visibly roughened or attacked

T = Thicker after etch (due to swelling or compound formation)

P = Some of film peeled during etch or when rinsed

I = Incubation time before etching fully starts

C = Film coagulated

soft = a soft material remained after etching

Evap = evaporated

Ion-Mill = ion-mill deposition

PECVD = plasma-enhanced chemical-vapor deposition

Si-Rich Si Nit. = silicon-rich silicon nitride

Stoich Si Nit. = stoichiometric silicon nitride

similar Borofloat glass) are used in anodic bonding to silicon due to the high content of mobile sodium ions and to the good match of thermal expansion rates. The large amounts of non-silicon-dioxide "impurities" give it noticeably different etching characteristics, etching slower in 5:1 BHF, but faster in silicon isotropic etchant.

*Thermal Oxide Wet-Grown:* Silicon dioxide grown in a Tylan atmospheric-pressure furnace with the recipe O<sub>2</sub> carrier

gas at 200 sccm, H<sub>2</sub>O vapor at a pressure just below 1 atm (the water source is at 98 °C) at 1100 °C, and a total pressure of 1 atm, followed by a 20-min N<sub>2</sub> anneal at 1100 °C. RI = 1.46. Thermal oxide forms a conformal coating on silicon. It is denser and etches more slowly than chemical-vapor-deposited oxides.

*Ann. LTO LPCVD Calogic:* Low-temperature silicon dioxide (LTO) deposited in a Calogic low-temperature

TABLE V  
 ETCH RATES OF AL, Ti, V, Nb, Ta, AND Cr (nm/min)

Etch	Aluminum Evap	Al + 2% Si Sputtered	Titanium Sputtered	Vanadium Evap	Niobium Ion-Mill	Tantalum Evap	Tantalum Ion-Mill	Chromium Evap	Chromium Ion-Mill	Plated Cr on Au Evap
Si Iso Etch	60	400	300	9600	79	5.8	5.3	R 8.8	-	< 2.9
KOH	12,900	F	soft	< 12	3.2	S	2.8	4.2	-	0
10:1 HF	W	250	1160	S	S	S	S	S	S	-
1:1 BHF	11	140	W	< 2	0	S	R 0	0	< 0.3	P
Pae Etch 4	1.9	R < 15	< 2	S	S	S	S	S	S	-
Phosphoric	> 500	980	-	-	0	-	0	100	-	-
Al Etch A	530	660	0	-	-	-	-	T 0	-	1.0
Ti Etch	150	240	1100	-	-	-	-	0	S	-
CR-7	3.8	S	< 2	60	R 0	S	< 0.7	170	150	110
CR-14	0	0.8	< 2	15	-	-	-	90	W	120
Mix Etch	> 20	-	-	-	-	-	-	R 0	-	-
H <sub>2</sub> O <sub>2</sub> 50°C	T 0	0.25	-	-	-	-	-	110	W	-
Cu FeCl <sub>3</sub> 200	35	W	-	-	-	-	-	0.053	S	-
Cu APS 100	< 0.3	-	-	-	-	-	-	0	S	-
Dil. Aqua regia	600	W	< 0.5	-	0	S	< 2	0	S	-
Alu-5	-	-	-	-	-	-	-	0	S	-
NiCr TFN	> 46	-	-	-	-	-	-	> 170	W	W
Phos-Sulf	W	W	-	-	-	-	-	I > 500	-	-
Piranha	> 5200	W	240	-	6.3	S	T 0	> 16	5.7	R 0
Microstrip	-	-	-	-	-	-	-	-	-	-
Acetone	S	0	0	S	S	S	S	S	S	S
Methanol	S	S	S	S	S	S	S	S	S	S
IPA	S	S	S	S	S	S	S	S	S	S
XeF <sub>2</sub>	S	0	29	W	W	W	W	-	-	-
HF vapor	R	R	R	-	-	-	-	S	S	-
Technics O <sub>2</sub>	S	0	0	S	S	S	S	S	S	S
DRIE HF mech.	-	-	4.8	-	-	-	-	-	-	-
DRIE HF ES	-	-	-	-	-	-	-	-	-	-
DRIE LF mech.	-	-	-	-	-	-	-	-	-	-
DRIE LF ES	-	-	-	-	-	-	-	-	-	-
STS 320 SF <sub>6</sub>	-	-	-	-	W	W	W	< 1	< 0.7	S
STS SF <sub>6</sub> +O <sub>2</sub>	< 2.8	-	-	-	26	W	37	< 1	< 0.8	S
STS 320 CF <sub>4</sub>	S	S	-	-	-	-	-	< 1	< 0.9	-
STS CF <sub>4</sub> +O <sub>2</sub>	0.87	1.5	-	-	14	-	21	< 1.3	< 1.2	-
Ion Mill	73	W	38	W	W	42	42	58	58	W

Notation:

 W = Etch known to work, but etch rate not measured  
 F = Etch known to be fast, but etch rate not measured  
 S = Etch rate known to be slow or zero, but etch rate not measured  
 R = Film was visibly roughened or attacked  
 T = Thicker after etch (due to swelling or compound formation)  
 P = Some of film peeled during etch or when rinsed  
 I = Incubation time before etching fully starts  
 C = Film coagulated  
 soft = a soft material remained after etching

 Evap = evaporated  
 Ion-Mill = Ion-mill deposited

chemical-vapor-deposition (LPCVD) furnace with the recipe  $\text{SiH}_4 = 74 \text{ sccm}$ ,  $\text{O}_2 = 105 \text{ sccm}$ ,  $p = 500 \text{ mtorr}$ ,  $T = 495^\circ\text{C}$ ,  $\text{RI} = 1.46$ , then annealed in  $\text{N}_2$  at  $1000^\circ\text{C}$  for 60 min. LTO is used instead of thermal oxide in many applications due to its relatively low deposition temperature and the ability to deposit thick films on surfaces other than silicon. LTO should be annealed prior to the deposition of another film at higher

temperature (e.g., polysilicon) because it will outgas hydrogen, which can cause the overlying film to bubble.

**Unan. LTO LPCVD Tylan:** LTO deposited in a Tylan LPCVD furnace with the recipe  $\text{SiH}_4 = 60 \text{ sccm}$ ,  $\text{O}_2 = 90 \text{ sccm}$ ,  $p = 300 \text{ mtorr}$ ,  $T = 450^\circ\text{C}$ , no anneal.

**Ann. LTO LPCVD Tylan:** LTO deposited as above, then annealed in  $\text{N}_2$  at  $1000^\circ\text{C}$  for 60 min.  $\text{RI} = 1.46$ .

TABLE VI  
 ETCH RATES OF Mo, W, Ni, Pd, Pt, Cu, Ag, Au, TiW, NiCr, AND TiN (nm/min)

Etch	Molybdenum Evap	Tungsten Sputtered	Nickel Evap	Palladium Evap	Platinum Evap	Copper Evap	Silver Evap	Gold Evap	10Ti/90W Ion-Mil	80Ni/20Cr Evap	TiN Sputtered
Si iso Etch	11,000	13	21	0	0	37	46	0	20	-	soft
KOH	0	0	0	0	0	T	0	T	0	> 300	-
10:1 HF	S	0	S	S	S	S	S	S	S	S	S
5:1 BHF	< 0.3	< 2	< 1.1	0	0	R	< 5	R	< 5	0	2.5
Pad Etch 4	S	S	-	-	-	-	-	-	-	-	-
Phosphoric	-	-	-	-	-	-	-	0	2.5	-	-
Al Etch A	-	-	29	-	-	> 2900	-	-	-	-	-
Ti Etch	-	11	-	-	-	-	-	-	0	-	-
CR-7	33	3.2	1.7	0	0	280	450	0	0.62	11.2	-
CR-14	soft	0	< 2	0	< 3	19	-	-	-	0.22	< 2
Moly Etch	850	-	-	-	-	-	-	-	0	-	-
H <sub>2</sub> O <sub>2</sub> 50°C	-	150	-	-	-	-	-	-	W	-	-
Cu FeCl <sub>2</sub> 200	-	-	21	-	-	3900	T	0	-	-	-
Cu APS 100	-	-	0	-	-	2500	-	-	-	-	-
Dil. Aqua regia	650	5.2	100	390	3.8	600	W	680	3.7	-	-
Al-5	-	-	0	T	0	T	0	650	-	-	-
NiCr TiN	680	-	10	-	-	690	-	-	-	83	-
Phos-Sulf	-	-	-	-	-	-	-	-	-	-	-
Piranha	18	-	380	3.0	< 3	86	600	0	0.78	92	-
Microstrip	-	-	-	-	-	-	-	-	-	-	-
Acetone	S	S	S	S	S	S	S	S	S	S	S
Methanol	S	S	S	S	S	S	S	S	S	S	S
IPA	S	S	S	S	S	S	S	S	S	S	S
XeF <sub>2</sub>	W	80	-	-	-	-	-	-	W	-	-
HF vapor	0	0	S	-	S	R	-	S	-	-	-
Technics O <sub>2</sub>	S	0	S	S	S	S	S	S	S	S	S
DRIE HF mech.	5.7	4.9	S	S	S	S	S	S	4.6	S	-
DRIE HF ES	-	-	-	-	-	-	-	-	-	-	-
DRIE LF mech.	-	-	-	-	-	-	-	-	-	-	-
DRIE LF ES	-	-	-	-	-	-	-	-	-	-	-
STS 320 SF <sub>6</sub>	73	W	S	S	S	S	-	S	W	S	-
STS SF <sub>6</sub> /O <sub>2</sub>	120	W	0.71	3.1	7.4	S	-	S	550	3.7	-
STS 320 CF <sub>4</sub>	W	-	-	-	-	-	-	S	-	-	-
STS CF <sub>4</sub> /O <sub>2</sub>	150	-	-	1.0	1.4	-	-	8.3	49	0	-
Ion Mill	54	38	66	130	88	110	220	170	W	18	W

Notation:

 W = Etch known to work, but etch rate not measured  
 F = Etch known to be fast, but etch rate not measured  
 S = Etch rate known to be slow or zero, but etch rate not measured  
 R = Film was visibly roughened or attacked  
 T = Thicker after etch (due to swelling or compound formation)  
 P = Some of film peeled during etch or when rinsed  
 I = Incubation time before etching fully starts  
 C = Film coagulated  
 soft = a soft material remained after etching

Evap = evaporated

Ion-Mil = Ion-mil deposited

**Unan. PSG LPCVD Tylan:** Phosphosilicate glass (PSG) deposited in a Tylan LPCVD furnace with the recipe  $\text{SiH}_4 = 60$  sccm,  $\text{O}_2 = 90$  sccm,  $\text{PH}_3 = 10.3$  sccm,  $p = 300$  mtorr,  $T = 450^\circ\text{C}$ , no anneal,  $\text{RI} = 1.47$ . PSG is LTO doped with  $\text{P}_2\text{O}_5$ , typically in the 5–10% range. It is used as a sacrificial layer in micromachining because it etches much faster than undoped LTO in HF solutions.

**Ann. PSG LPCVD Tylan:** PSG deposited as above, then annealed in  $\text{N}_2$  at  $1000^\circ\text{C}$  for 60 min.  $\text{RI} = 1.48$ . Like pure LTO, PSG should be annealed prior to the deposition of another film

at high temperature (e.g., polysilicon) because it will outgas hydrogen, which can cause the overlying film to bubble. Annealing results in a lower etch rate in HF solutions.

**Oxide PECVD Unannealed:** Silicon dioxide deposited in an STS 310 plasma-enhanced chemical-vapor-deposition (PECVD) system with the recipe  $2\%\text{SiH}_4/\text{Ar} = 500$  sccm,  $\text{N}_2\text{O} = 1110$  sccm,  $P = 10$  W at  $13.56$  MHz,  $p = 800$  mtorr,  $T = 300^\circ\text{C}$ ,  $\text{RI} = 1.46$ . PECVD silicon dioxide usually contains 2–9 atomic % hydrogen [5]. PECVD oxide can be deposited at an even lower temperature than LPCVD films.

TABLE VII  
 ETCH RATES OF RESISTS, PARYLENE, AND POLYIMIDE (nm/min)

Etch	S1822 Positive Photoresist	OOG 820 Positive Photoresist	Futurrex Positive Photoresist	Futurrex Negative Photoresist	Act. Mkr. Resist Pen	Parylene Type C	Spin-On PI 2555 Polyimide
Si Iso Etch	P 0	0	P 21	290	-	0.019	0
KOH	> 17,600	F	> 13,000	> 18,000	F	0.42	T 0
10:1 HF	S	0	-	-	S	4.4	S
5:1 BHF	0	0	21	5.0	< 3	0.16	5.5
Pad Etch 4	T 0	-	-	-	-	-	-
Phosphoric	P 120	55	P 77	> 400	-	0.55	-
Al Etch A	0	0	-	-	-	-	-
Ti Etch	< 0.5	0	-	-	-	-	-
CR-7	0	S	0	< 0.5	38	-	0
CR-14	0.24	S	S	S	S	-	S
Moly Etch	230	-	-	-	-	-	-
H <sub>2</sub> O <sub>2</sub> 50°C	R T 0	S	S	S	S	T 0	S
Cu FeCl <sub>3</sub> 200	0.48	S	S	S	S	-	S
Cu APS 100	0	S	S	S	S	-	S
Dil. Aqua regia	0	S	T 0	T 0	T 0	-	T 0
AU-5	P 0	S	S	S	S	-	S
NiCr TFN	0.57	S	S	S	S	-	S
Phos+Sulf	F	F	F	F	F	-	-
Piranha	> 92,000	F	> 50,000	> 50,000	> 15,000	2.6	> 17,000
Microstrip	> 94,000	F	> 85,000	> 90,000	> 11,000	-	520
Acetone	> 178,000	F	> 120,000	> 87,000	> 26,000	0.77	0
Methanol	> 35,000	-	P C > 16,000	> 27,000	P	0.093	0
IPA	> 1000	-	480	910	-	T 0	0
XaF <sub>2</sub>	S	0	S	S	S	-	S
HF vapor	-	P 0	S	S	S	-	S
Technics O <sub>2</sub>	300	340	200	470	370	220	370
DRIE HF mech.	30	W	W	W	54	-	-
DRIE HF ES	W	W	W	W	W	27	-
DRIE LF mech.	35	W	34	W	W	-	-
DRIE LF ES	W	W	W	W	W	15	-
STS 320 SF <sub>6</sub>	120	W	I 64	140	W	-	140
STS SF <sub>6</sub> +O <sub>2</sub>	100	-	I 130	320	170	-	200
STS 320 CF <sub>4</sub>	42	W	W	72	W	-	-
STS CF <sub>4</sub> +O <sub>2</sub>	130	-	I 100	190	52	-	160
Ion Mill	R < 100	W	W	W	W	W	W

Notation:

W = Etch known to work, but etch rate not measured  
 F = Etch known to be fast, but etch rate not measured  
 S = Etch rate known to be slow or zero, but etch rate not measured  
 R = Film was visibly roughened or attacked  
 T = Thicker after etch (due to swelling or compound formation)  
 P = Some of film peeling during etch or when rinsed  
 I = Incubation time before etching fully starts  
 C = Film congealed  
 soft = a soft material remained after etching

Act. Mkr. = Action Marker

*Oxide PECVD Annealed:* Silicon dioxide deposited as above, then annealed in N<sub>2</sub> at 1000 °C for 60 min. RI = 1.46. Annealing densifies the film, resulting in a lower etch in HF solutions.

*Oxide Ion-Milled:* Silicon dioxide ion-mill-deposited from a silicon-dioxide target in a Commonwealth Scientific system with argon ions at 1250 V, current density of about 2 mA/cm<sup>2</sup>, p = 3 × 10<sup>-4</sup> torr. RI = 1.57. Ion milling is the lowest temper-

ature deposition method studied here, with the wafer remaining near room temperature.

### C. Silicon Nitride

*Stoich. Si Nit. LPCVD:* Stoichiometric silicon nitride (Si<sub>3</sub>N<sub>4</sub>) deposited in a Tylan LPCVD furnace with the recipe SiCl<sub>2</sub>H<sub>2</sub> = 25 sccm, NH<sub>3</sub> = 75 sccm, p = 200 mtorr, T = 800 °C, RI = 1.99. Silicon nitride is used as an etch

mask, thermal-oxidation mask, and wafer-capping material due to its low permeability to moisture. It typically has a high tensile residual stress.

**Si-Rich Si Nit. LPCVD:** Silicon-rich, low-stress silicon nitride deposited in a Calogic LPCVD furnace with the recipe  $\text{SiCl}_2\text{H}_2 = 159 \text{ sccm}$ ,  $\text{NH}_3 = 40.5 \text{ sccm}$ ,  $p = 280 \text{ mtorr}$ ,  $T = 850^\circ\text{C}$ ,  $\text{RI} = 2.24$ . Low-stress silicon nitride typically has a tensile residual stress below 50 MPa, allowing its use in micromachining as a freestanding film.

**PECVD Silicon Nit. Low RI:** Silicon nitride deposited in an STS 310 PECVD system with a relatively low ratio of silicon to nitrogen with the recipe  $2\%\text{SiH}_4/\text{Ar} = 100 \text{ sccm}$ ,  $\text{N}_2 = 2000 \text{ sccm}$ , power alternating between  $P = 60 \text{ W}$  at  $13.56 \text{ MHz}$  for 6 s and  $25 \text{ W}$  at  $187 \text{ kHz}$  for 3 s,  $p = 750 \text{ mtorr}$ ,  $T = 350^\circ\text{C}$ ,  $\text{RI} = 1.88$ . PECVD silicon nitride is deposited at a much lower temperature than LPCVD and can contain well over 10 atomic percent hydrogen [5], and is therefore frequently written as  $\text{Si}_3\text{N}_4\text{H}_x$  or  $\text{SiN}_x\text{H}_y$ . High frequency typically yields tensile silicon nitride films; low frequency typically gives compressive films. "Mixed-frequency" nitride such as this has a lower absolute value of stress.

**PECVD Silicon Nit. High RI:** Silicon nitride deposited as above, but with a relatively high ratio of silicon to nitrogen by increasing the flow of  $2\%\text{SiH}_4/\text{Ar}$  to 500 sccm.  $\text{RI} = 2.53$ . This PECVD silicon nitride has a significantly lower etch rate in HF solutions than its low-index counterpart above, but still etches much faster than LPCVD nitride.

## D. Aluminum Oxide

**Sapphire Wafer:** Clear, colorless, pure crystalline aluminum oxide wafer. (Aluminum oxide is also known as alumina or corundum.) These wafers are referred to as sapphire in the electronics industry, although true sapphire has a small amount of titanium, giving it a blue color. Sapphire has been used as substrate for silicon-on-insulator (SOI) wafers. Sapphire was the most chemically inert material tested for this paper.

**Aluminum Oxide Ion-Milled:** Aluminum oxide ion-milled deposited in a Commonwealth Scientific system from a sapphire target with argon ions at 1250 V, current density of about  $2 \text{ mA/cm}^2$ . The sapphire was held to a copper backing plate with copper-loaded vacuum grease. A small area of this grease was exposed during deposition. Adhesion of the film to the wafer was good. Like silicon dioxide, aluminum oxide is relatively inert. Aluminum oxide films have been used as dielectrics and protective coatings.

**Aluminum Oxide Evaporated:** Aluminum oxide electron-beam evaporated from an alumina source. Wafer temperature was not controlled and was near room temperature during deposition.  $\text{RI} = 1.66$ .

## E. Aluminum

**Aluminum Evaporated:** Aluminum evaporated from a tungsten wire in a Veeco evaporator. Aluminum is commonly used for electrical interconnections due to its ease of use and good conductivity. It is well known to have good adhesion to silicon and silicon dioxide.

**Al + 2% Si Sputtered:** Aluminum sputtered using argon in a CPA 9900 sputtering system with the recipe  $p = 6 \text{ mtorr}$ ,

$P = 4.5 \text{ kW}$ , track speed =  $10 \text{ cm/min}$ . Aluminum with a small percentage of silicon, added to prevent "spiking" of metal into the substrate, is a common material for IC interconnects.

## F. Transition Metals

**Titanium Sputtered:** Titanium sputtered using argon in a CPA 9900 sputtering system with the recipe  $p = 6 \text{ mtorr}$ ,  $P = 4.5 \text{ kW}$ , track speed =  $10 \text{ cm/min}$ . Titanium is well known to have good adhesion to silicon and silicon dioxide, and is frequently used as an adhesion layer for less-adhesive metals such as gold.

**Vanadium Evaporated:** Vanadium electron-beam evaporated in a CHA evaporator at a rate of  $0.1 \text{ nm/s}$ . The pressure was below  $9 \times 10^{-7} \text{ torr}$  at the start of the evaporation. Vanadium was found to have good adhesion to silicon and silicon dioxide, as evaluated by firmly pressing 3M Scotch Tape onto the film, then peeling the tape off without removing the metal. Vanadium thus has potential as an alternative adhesion layer.

**Niobium Ion Milled:** Niobium ion-mill-deposited in a Commonwealth Scientific system with argon ions at 1250 V, current density of about  $2 \text{ mA/cm}^2$ ,  $p = 3 \times 10^{-4} \text{ torr}$ . Niobium was found to have good adhesion to silicon, so it has potential as an alternative adhesion layer.

**Tantalum Evaporated:** Tantalum electron-beam evaporated in a CHA evaporator at a rate of  $0.1 \text{ nm/s}$ . The pressure was below  $9 \times 10^{-7} \text{ torr}$  at the start of the evaporation. Tantalum was found to have good adhesion to silicon and silicon dioxide, so it has potential as an alternative adhesion layer. As one of the high-melting-temperature metals, tantalum has been used as a hot filament in inert atmospheres.

**Tantalum Ion-Milled:** Tantalum ion-mill-deposited in a Commonwealth Scientific system with argon ions at 1250 V, current density of about  $2 \text{ mA/cm}^2$ ,  $p = 3 \times 10^{-4} \text{ torr}$ . Tantalum deposited using this method also was found to have good adhesion to silicon and silicon dioxide.

**Chromium Evaporated:** Chromium evaporated from a tungsten strip in a CHA evaporator at a rate of  $0.1 \text{ nm/s}$ . The pressure was below  $9 \times 10^{-7} \text{ torr}$  at the start of the evaporation. Chromium is well known to have good adhesion to silicon and silicon dioxide, and is frequently used as an adhesion layer for less-adhesive metals such as gold.

**Chromium Ion Milled:** Chromium ion-mill-deposited in a Commonwealth Scientific system with argon ions at 1250 V, current density of about  $2 \text{ mA/cm}^2$ ,  $p = 3 \times 10^{-4} \text{ torr}$ .

**Patterned Cr on Au Evaporated:** Evaporated chromium previously lithographically patterned and etched, sitting on a continuous layer of gold (which itself has a chromium adhesion layer). This was done to study the potential electrochemical effect of this commonly used system of two metals in contact.

**Moly Evaporated:** Molybdenum electron-beam evaporated in a CHA evaporator at a rate of  $0.5 \text{ nm/s}$ . The pressure was below  $9 \times 10^{-7} \text{ torr}$  at the start of the evaporation. The evaporated molybdenum used in these tests had good adhesion to silicon, but was found to have only fair adhesion to oxide; it did not peel when Scotch tape was lightly pressed, but did when firmly pressed and removed. We have found it to have good adhesion when sputtered. Molybdenum has a relatively low thermal coefficient of expansion and has been used as a heat sink attached

to silicon. It has been used in MEMS as the electrode material for film-bulk acoustic resonators [6].

**Tungsten Sputtered:** Tungsten sputtered using argon in a CPA 9900 sputtering system with the recipe  $p = 6$  mtorr,  $P = 1.5$  kW, track speed =  $10$  cm/min. An adhesion layer of titanium was used under the tungsten. Tungsten also has a relatively low thermal coefficient of expansion and has been used as a heat sink attached to silicon. Tungsten has been used in MEMS as a hot filament in vacuum [7] and as an interconnect material compatible with high-temperature processing.

**Nickel Evaporated:** Nickel electron-beam evaporated in a CHA evaporator at a rate of  $0.3$  nm/s. The pressure was below  $9 \times 10^{-7}$  torr at the start of the evaporation. A  $10$ -nm adhesion layer of chromium was used under the nickel. Nickel has been used as a diffusion-barrier layer in contacts, a seed layer for electroplating (immediately after removing its native oxide), and as the metal to which solder adheres.

**Palladium Evaporated:** Palladium electron-beam evaporated in a CHA evaporator at a rate of  $0.15$  nm/s. The pressure was below  $9 \times 10^{-7}$  torr at the start of the evaporation. A  $10$ -nm adhesion layer of chromium was used under the palladium. Like platinum, palladium has been used as catalyst, for example in the hydrogenation of carbon compounds [2]. In these tests, it was etched more easily than platinum.

**Platinum Evaporated:** Platinum electron-beam evaporated in a CHA evaporator at a rate of  $0.15$  nm/s. The pressure was below  $9 \times 10^{-7}$  torr at the start of the evaporation. A  $10$ -nm adhesion layer of chromium was used under the platinum. Platinum has been used in thin-film heaters, resistive temperature sensors, and as a diffusion-barrier layer in contacts due to its inertness and stability. It is also well-known as a catalyst and has been used in fine-grain form (for large surface area) in microscale fuel cells. Platinum is often used as the electrode layer on which lead zirconate titanate (PZT) is grown for good crystal orientation. In this work it was the most difficult metal to etch.

**Copper Evaporated:** Copper evaporated from a tungsten boat in a CHA evaporator at a rate of  $0.17$  nm/s. The pressure was below  $9 \times 10^{-7}$  torr at the start of the evaporation. A  $10$ -nm adhesion layer of chromium was used under the copper. Copper is second to only to silver in its electrical conductivity, and is presently used in high-speed integrated circuits. Plated copper has been employed in MEMS for high-Q inductors.

**Silver Evaporated:** Silver electron-beam evaporated in a CHA evaporator at a rate of  $0.5$  nm/s. The pressure was below  $9 \times 10^{-7}$  torr at the start of the evaporation. A  $10$ -nm adhesion layer of chromium was used under the silver. Silver was found to form a dark tarnish (due to sulfide formation, not oxidation [8]) within days of deposition.

**Gold Evaporated:** Gold electron-beam evaporated in a CHA evaporator at a rate of  $0.5$  nm/s. The pressure was below  $9 \times 10^{-7}$  torr at the start of the evaporation. A  $10$ -nm adhesion layer of chromium was used under the gold. Gold, which is an excellent electrical conductor, has many applications in electronics: Because it does not corrode like copper and aluminum, it has been used in mechanical contacts and bond pads. Micromachined switches have employed gold or its alloys. Gold surfaces are wetted well by many solders, but are also rapidly dissolved, so an underlying metal layer (e.g., Ni or Pt) must be used.

## G. Metal Alloys and Compound

**10 Ti/90 W Ion-Milled:** 10% titanium/90% tungsten alloy ion-mill-deposited in a Commonwealth Scientific system with argon ions at  $1250$  V, current density of about  $2$  mA/cm<sup>2</sup>,  $p = 3 \times 10^{-1}$  torr. This alloy is commonly used as a combination adhesion/diffusion-barrier layer.

**80 Ni/20 Cr Evaporated:** 80% nickel/20% chromium (at the source) alloy evaporated from a tungsten wire in a Veeco evaporator. Due to their inertness at higher temperature and relatively low thermal conductivity, nickel-chromium alloys have been used for heaters.

**TiN Sputtered:** Titanium nitride sputtered using in a Varian/Novellus M12 sputtering system with the recipe argon =  $55$  sccm, nitrogen =  $65$  sccm,  $P = 7.2$  kW,  $T = 375$  °C. TiN is used as an adhesion layer.

## H. Resists

**S1822 Positive Photoresist:** Shipley Microposit S1822 positive photoresist, dispensed, and spun at  $4600$  rpm for  $30$  s for a thickness of  $2.2$   $\mu$ m, soft baked on a hot plate at  $115$  °C for  $60$  s, then hard baked in an oven at  $110$  °C for  $30$  min RI =  $1.65$ .

**OCG Positive 820 Photoresist:** OCG (formerly KTI) 820 photoresist, dispensed, and spun at  $6000$  rpm for  $30$  s for a thickness of  $1.1$   $\mu$ m, soft baked on a hot plate, then hard baked in an oven at  $120$  °C for  $30$  min RI =  $1.65$ .

**Futurrex Positive Photoresist:** Futurrex PR1-4000A photoresist, dispensed, and spun at  $2000$  rpm for  $45$  s for a thickness of  $4$   $\mu$ m, and soft baked at  $120$  °C for  $90$  s on a hot plate (no hard bake was done per the manufacturer's instructions). RI =  $1.65$ . This photoresist is formulated for a slow plasma-etch rate. In SF<sub>6</sub> and CF<sub>4</sub>-based plasmas, it was found to barely etch for about  $10$  min (an "incubation time"), after which it started to etch more rapidly.

**Futurrex Negative Photoresist:** Futurrex NR5-8000 photoresist, dispensed, and spun at  $5000$  rpm for  $45$  s for a thickness of  $6$   $\mu$ m, and soft baked at  $145$  °C for  $60$  s on a hot plate (no hard bake was done per the manufacturer's instructions). Unlike most photoresists, Futurrex positive and negative photoresists use the same developer.

**Action Marker Resist Pen:** ITW Dymon Action Marker Fine Line 33 PC-901 Blue resist pen, written onto a wafer and allowed to dry (no bake). This resist pen is convenient for rapidly masking an area or touching up photoresist before a wet or dry etch. It is rapidly stripped in acetone or photoresist stripper. Thicknesses were measured using profilometry.

## I. Parylene and Polyimide

**Parylene Type C:** Parylene Type C deposited using a Specialty Coating Systems Model PDS 2010 Labcoat 2 system by heating the solid source material, the dimer di-para-xylylene, to  $175$  °C to sublimate it, then heating the dimer vapor to  $690$  °C to cleave the molecules into the monomer para-xylylene, finally depositing it on the substrate at room temperature and  $p = 25$  mtorr to form a film of the polymer poly-(para-xylylene). Parylene is the trade name of the polymer, given by Union Carbide. RI =  $1.65$ .

There are several types of Parylene: Parylene-N can be envisioned as a polymer of benzene rings with  $\text{CH}_2$  groups on opposite ends. Parylene-C has a chlorine atom substituted for one of the hydrogen atoms in the benzene rings, while Parylene-D has two such substitutions. Parylenes are very inert chemically: of the etches tested, Parylene-C only had a significant etch rate in the oxygen plasma and ion mill (it also etched in a separate test in an  $\text{SF}_6$  plasma). Deposition is conformal and pinhole-free. Parylenes are also biocompatible and have a relatively low gas permeability. The flexibility of Parylene films has allowed their use in such MEMS applications as miniature valves and wings [9].

**Spin-On PI 2556 Polyimide:** DuPont PI2556 spin-on polyimide, dispensed, and spun at 4000 rpm for 60 s for a thickness of 1.1  $\mu\text{m}$ , held at 20 °C for 30 min, then baked at 100 °C for 30 min, and finally at 250 °C for 120 min (the manufacturer's instructions call for the last bake to be 300 °C for complete crosslinking). RI = 1.82.

### III. ETCH DESCRIPTIONS AND INFORMATION

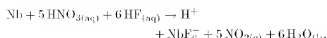
The wet and dry etches are described below, listed by the abbreviation labels (in italics) in the etch-rate tables. Information about the etch properties is included. Further documentation, including many chemical reactions, was presented previously [3].

#### A. Wet Etches

***Si Iso Etch:*** Isotropic Silicon Etchant "Trilogy Etch," mixed from 126 parts  $\text{HNO}_3$  : 60 parts  $\text{H}_2\text{O}$  : 5 parts  $\text{NH}_4\text{F}$ .  $T \approx 20^\circ\text{C}$  (room temperature). This solution should be mixed several hours before use to yield a stable etch rate. Similar commercial preparations are also available. During use, agitation increases uniformity. The solution gets weaker with use. Silicon etching in hydrofluoric acid-nitric acid-water (HF-nitric) and hydrofluoric acid-nitric acid-acetic acid (HNA) etch systems has been well characterized by Schwartz and Robbins [10].

This etchant is intended for isotropically etching silicon, both polycrystalline and single-crystal. It can be used to round off sharp corners formed during orientation-dependent etching and scallops and ridges formed during deep-reactive ion etching (DRIE) to lower the stress concentration of mechanical components. Silicon iso etch was also found to etch graphite, vanadium, niobium, and molybdenum at moderate to very high rates. Due to the small amount of HF formed upon mixing, it also slowly etches silicon dioxide. Photoresist can be normally be used as a mask with this etchant, although two of the resists used in these tests peeled.

Silicon and many metals are etched by the same sequence of steps in HNA etchants: Nitric acid is a powerful oxidizing agent. In nitric acid alone, the silicon or metal is oxidized to form a tough oxide film. The addition of hydrofluoric acid to the solution causes some of these metal oxides to be etched. A representative overall reaction is that for niobium [11]:



***KOH:*** 30% potassium hydroxide (by weight) in water.  $T = 80^\circ\text{C}$ . This solution can be mixed from solid pellets and water,

TABLE VIII  
ETCH RATES OF THERMAL OXIDE IN VARIOUS DILUTIONS OF HF AND BHF (nm/min)

Etchant	Thermal Oxide Etch Rate
Conc. HF (49%)	2300
10:1 HF	23
25:1 HF	9.7
100:1 HF	2.3
5:1 BHF	100
10:1 BHF	50

Notation:  
BHF = buffered hydrofluoric acid  
conc. = concentrated

or from a diluted commercially prepared aqueous solution. It is used for crystal-orientation-dependent etching (ODE) of silicon, for etching n-type silicon to a heavily-doped p-type etch stop, and for electrochemically etching p-type silicon to an n-type etch stop. Data on etch rates at different KOH concentrations and temperatures and different silicon doping levels, as well as etch-stop mechanisms, have been documented by Seidel *et al.* [12], [13]. KOH and other orientation-dependent etches have been summarized by Williams [14].

While KOH is most often used for ODE of single-crystal silicon (SCS), it can also be used for rapid etching of shallow cavities of any shape, if the exact shape at the perimeter is not of concern. KOH has a high etch rate for both polycrystalline silicon and germanium, as well as for heavily ion-implanted SCS that has not been annealed.

KOH was found to etch silicon dioxides, LPCVD silicon nitride, and most of the transition metals tested slowly. High-refractive index PECVD nitride etched sufficiently slowly to be useful as a mask. Aluminum and photoresist were very rapidly etched. KOH is commonly masked with LPCVD silicon nitride, a sandwich of PECVD oxide and nitride (to reduce pinholes), or gold with a chromium adhesion layer.

***10:1 HF:*** Mixed from 10 parts  $\text{H}_2\text{O}$ :1 part 49% HF, by volume.  $T \approx 20^\circ\text{C}$  (room temperature). HF etches silicon dioxide. Such diluted HF solutions are used to strip thin layers of oxide and native oxide. The solution gets weaker with use. The etch rates of thermal oxide in various dilutions of HF are given in Table VIII. The etch rate is linear with concentration for dilutions in the 10:1 to 100:1 range.

***5:1 BHF:*** Mixed from 5 parts 40%  $\text{NH}_4\text{F}$ :1 part 49% HF, by volume.  $T \approx 20^\circ\text{C}$  (room temperature). After mixing, buffered hydrofluoric acid (BHF) solutions should sit for several hours to stabilize, as the etch rate will initially be much higher than in equilibrium. Commercial mixes are also available. BHF, also known as buffered oxide etch (BOE), is used for its stable etch rate with use. The etch rates of thermal oxide in 5:1 and 10:1 BHF are also given in Table VIII. The etch rate in 10:1 BHF is exactly half that in 5:1 BHF.

***Pad Etch 4:*** Pad Etch 4 from Ashland is a commercial mix of 11–15%  $\text{NH}_4\text{F}$  + 30–34%  $\text{CH}_3\text{COOH}$  + 47–51%  $\text{H}_2\text{O}$  + 4–8% propylene glycol + surfactant [15].  $T \approx 20^\circ\text{C}$  (room tempera-



ture). This etchant is formulated to etch through a silicon dioxide dielectric layer, with the etch slowing greatly on aluminum pads or contacts. In these tests pure Al etched only slowly, but a long exposure was found to roughen the surface of Al/2%Si, making it impossible to report an etch rate. Photoresist can be used as an etch mask for Pad Etch.

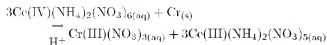
**Phosphoric:** Phosphoric Acid (85% by weight), 160 °C. It is one of the few wet etches for silicon nitride. Because it etches photoresist, another masking material, such as oxide, must be used. The etch rates reported here for stoichiometric and silicon-rich LPCVD silicon nitrides are faster than reported earlier [3]. The etch rate is very sensitive to temperature, and this bath was likely a few degrees hotter. Hot phosphoric acid was also found to rapidly etch aluminum.

**Al Etch A:** Aluminum Etchant Type A from Transene is a commercial mix of 80%  $\text{H}_3\text{PO}_4$  + 5%  $\text{HNO}_3$  + 5%  $\text{CH}_3\text{COOH}$  + 10%  $\text{H}_2\text{O}$  [16].  $T \approx 50^\circ\text{C}$ . It is formulated to selectively etch aluminum. It was also found to rapidly etch copper. Photoresist can be used as an etch mask for this etchant.

**Ti Etch:** Titanium wet etchant is mixed from 20 parts  $\text{H}_2\text{O}$ :1 part 30%  $\text{H}_2\text{O}_2$ :1 part 49% HF, by volume.  $T \approx 20^\circ\text{C}$  (room temperature). Being a dilute HF solution, this etches oxides relatively slowly. Photoresist can be used as an etch mask for this etchant.

**CR-7:** Chromium etchant CR-7 from Cyantek is a commercial mix of 9%  $(\text{NH}_4)_2\text{Ce}(\text{NO}_3)_6$  + 6%  $\text{HClO}_4$  +  $\text{H}_2\text{O}$  [17].  $T \approx 20^\circ\text{C}$  (room temperature). This etchant is formulated to selectively etch chromium, and was also found to etch copper and silver at useful rates. The manufacturer lists its shelf life as 12 months. Photoresist can be used as an etch mask for this etchant.

The chemical reaction for etching Cr is [18]



where the  $\text{H}^+$  on the arrow indicates an acidic solution. The  $\text{Cr}^{4+}$  ions are reduced to  $\text{Cr}^{3+}$ , while the Cr metal is oxidized to  $\text{Cr}^{3+}$ . The etch products are soluble, and further reactions may occur in solution [18]. The perchloric acid is a stabilizer for the ceric ammonium nitrate.

**CR-14:** Chromium etchant CR-14 from Cyantek is a commercial mix of 22%  $(\text{NH}_4)_2\text{Ce}(\text{NO}_3)_6$  + 8%  $\text{CH}_3\text{COOH}$  +  $\text{H}_2\text{O}$  [19].  $T \approx 20^\circ\text{C}$  (room temperature). This etchant is also formulated to selectively etch chromium. It was also found to slowly etch vanadium and copper. When molybdenum was etched, it left behind a soft compound. The manufacturer lists its shelf life as 3 months. It etches chromium a little slower than CR-7, but over several years of use we found it to have more a consistent etch rate, as well as much less undercut of photoresist. The etch reaction is the same as for CR-7. The stabilizer is instead acetic acid.

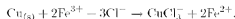
**Moly Etch:** Molybdenum wet etchant is mixed from 180 parts  $\text{H}_3\text{PO}_4$ :11 parts  $\text{CH}_3\text{COOH}$ :11 parts 70%  $\text{HNO}_3$ :50 parts  $\text{H}_2\text{O}$ , by volume.  $T \approx 20^\circ\text{C}$  (room temperature). It can be masked with photoresist, which is etched at a moderate rate.

$\text{H}_2\text{O}_2$  50 °C: Hydrogen peroxide (30% by weight).  $T = 50^\circ\text{C}$ . Hydrogen peroxide targets tungsten, but etches it slowly.

Heating increases the etch rate. Ultrasonic agitation (not done here) aids in etch uniformity by helping to remove bubbles. Warm  $\text{H}_2\text{O}_2$  was also found to slowly etch chromium.

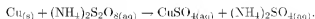
**Cu  $\text{FeCl}_4$  200:** Copper etchant type CE-200 from Transene is a commercial mix of 30%  $\text{FeCl}_3$  + 3–1%  $\text{HCl}$  +  $\text{H}_2\text{O}$  [20].  $T \approx 20^\circ\text{C}$  (room temperature). When silver was placed in the solution, the film became thicker. CE-200 can be patterned with photoresist.

$\text{FeCl}_3$  dissociates in water to generate  $\text{Fe}^{3+}$  and  $\text{Cl}^-$  ions. One of the reactions for etching of copper with ferric chloride is [21]



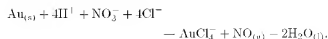
Other chloride complexes are also formed. The  $\text{Fe}^{3+}$  ions are reduced to  $\text{Fe}^{2+}$ , which remain in solution, while the copper metal is oxidized to  $\text{Cu}^{2+}$ .  $\text{HCl}$  assists in the dissolution of the ferric chloride, and also etches copper itself [21]. The manufacturer recommends rinsing in a 5–10%  $\text{HCl}$  solution after etching (not done here) before a final water rinse.

**Cu APS 100:** Copper etchant APS 100 from Transene is a commercial mix of 15–20%  $(\text{NH}_4)_2\text{S}_2\text{O}_8$  +  $\text{H}_2\text{O}$  [22].  $T = 30^\circ\text{C}$ . Photoresist can be used as an etch mask for this etchant. The overall reaction for ammonium persulfate etching of copper is [21]



**Dil. Aqua Regia:** A diluted solution of aqua regia mixed from 3 parts 37%  $\text{HCl}$ :1 part 70%  $\text{HNO}_3$ :2 parts  $\text{H}_2\text{O}$  (standard aqua regia does not have the additional water). The solution is self heating to about  $30^\circ\text{C}$  for tens of minutes for the shallow dish used in these tests. The water was added to the standard aqua regia solution to reduce the attack of photoresist. Aqua regia targets noble metals. In these tests, it was found to etch gold and palladium, but only slowly etch platinum (heating is require for a useful etch rate of Pt). It was also found to etch nickel, copper, molybdenum, and aluminum. Because the  $\text{HCl}$  evaporates off when left open, it should be mixed just before use.

Gold reacts with chlorine, bromine, and iodine at elevated temperatures to form the corresponding halides, but reacts only slowly with fluorine [23]. At room temperature, the nitric acid in aqua regia is an oxidizing agent, while the addition of hydrochloric acid provides nascent chlorine to form a soluble etch product. Similar overall reactions are given in the literature [8] and [23], including [24]



The  $\text{HCl}$  in aqua regia can be replaced with  $\text{HBr}$  or  $\text{HI}$  [23], but not  $\text{HF}$  [8], as gold does not form a soluble fluoride [25]. Thus,  $\text{HF}$ -nitric solutions such as the silicon isotropic etch do not etch gold.

**AU-5:** Gold etchant AU-5 from Cyantek is a commercial mix of 5%  $\text{I}_2$  + 10%  $\text{KI}$ –85%  $\text{H}_2\text{O}$  [26].  $T \approx 20^\circ\text{C}$  (room temperature). It is formulated to selectively etch gold. Such iodine solutions are the color of dark coffee, making it impossible to

observe a submerged sample. Rinsing with running water must be done to remove the solution from the surface. Isopropanol or methanol can be used for a faster rinse [18]. AU-5 can be patterned with photoresist. Rather than etching palladium, silver, and copper, it formed a compound that increased the film thickness above that of the original metal.

The overall reaction for etching gold is [18]



The potassium iodide increases the solubility of both the iodine and gold(I) iodide in water [18], [27], allowing a greater concentration of reactant in the solution and the etch product to be removed so that the etch can proceed.

**NiCr TFN:** Nichrome etchant TFN from Transene is a commercial mix of 10–20% (NH<sub>4</sub>)<sub>2</sub> Ce(NO<sub>3</sub>)<sub>6</sub> + 5–6% HNO<sub>3</sub>–H<sub>2</sub>O [28].  $T \approx 20^\circ\text{C}$  (room temperature). It is formulated to etch nickel-chromium alloy. It etched pure nickel slowly, but etched chromium, copper, and molybdenum faster than the NiCr alloy. Photoresist can be used as an etch mask for this etch.

**Phos + Sulf:** A mix of 1 part 96% H<sub>2</sub>SO<sub>4</sub>:1 part 85% H<sub>3</sub>PO<sub>4</sub>.  $T = 160^\circ\text{C}$ . This solution is listed as an etchant for sapphire [1], [2], but did not etch it even at this high temperature.

**Piranha:** The Piranha (also known as sulfuric-peroxide) used here is a mix of ~50 parts 96% H<sub>2</sub>SO<sub>4</sub>:1 part 30% H<sub>2</sub>O<sub>2</sub>.  $T = 120^\circ\text{C}$ . The hydrogen peroxide is added just before use. It is used as a cleaning solution that strips organics and some metals. In these tests, it was indeed found to etch photoresist, the resist pen, and polyimide very rapidly, but only slowly attacked the Parylene C. It also etched aluminum, nickel, and silver rapidly. Chromium was etched slowly, which allows it to be used for the cleaning of photomasks.

**Microstrip:** Arch Microstrip 2001 is a commercial photoresist-stripper mixed from 45–55% *n*-methyl pyrrolidone and 40–55% 2-(2-aminethoxy)ethanol [29].  $T = 85^\circ\text{C}$ . It is formulated to strip resists without affecting structural materials. In these tests, it very rapidly removed the photoresists and the resist pen, while etching the polyimide at a moderate rate and not etching graphite.

**Acetone:** Pure acetone, also known as dimethyl ketone. It is used to strip photoresist and for cleaning off organics. It removed photoresists rapidly, as expected, but did not etch polyimide.

**Methanol:** Pure methanol. It is used for cleaning and in supercritical drying. Like acetone, methanol removed photoresist, but did not etch polyimide.

**Isopropanol:** Pure isopropanol, also known as isopropyl alcohol (IPA) and 2-propanol. It is used for cleaning off organics. In contrast to methanol, isopropanol was found to remove photoresist at a moderate rate.

## B. Dry Etches

Most dry etches employ a plasma to generate reactive species, but several plasmaless-vapor-phase etches have been used in the fabrication of MEMS.

**XeF<sub>2</sub>, 2.6 mtorr, Homemade Chamber:** Xenon difluoride gas (no plasma) from a solid source at its full vapor pressure at room temperature. It targets silicon, as well as metals that form volatile fluorides such as Ti, W, Mo, and Ta [30]. It does not etch silicon dioxide, aluminum, or photoresist, and only slowly etches silicon nitride. Commercial XeF<sub>2</sub> etchers are now available. Other noble-gas fluorides have also been used to etch silicon [14].

**HF Vapor Over Dish:** HF+H<sub>2</sub>O vapor evaporated at room temperature from liquid 49% HF/H<sub>2</sub>O, with wafer continuously exposed ~1 cm over the dish. Like liquid HF, HF vapor targets silicon dioxides. Continuous exposure to the vapor results in droplet condensation on the wafer after about 25 s at room temperature, with the underlying areas having higher oxide etch rate. "Pulsed" etching, in which the wafer is alternately held over the liquid for 15 s, then removed and optionally heated for 15 s, eliminates droplet formation. Simply heating the wafer while over the HF/H<sub>2</sub>O also inhibits condensation, while still allowing etching to occur. In these tests, photoresist survived etching, but peeled when rinsed.

Manual vapor etching of oxide is particularly useful in cases where fragile structures cannot tolerate loading into a plasma etcher and where high selectivity is needed. Other users of this etch have found a nonvolatile residue left behind when doped oxides such as PSG [31] or silicon oxynitride [32] are exposed to HF vapor. Titanium and copper have been found to form non-volatile fluoride layers [31]. While we found aluminum to be attacked, it has also been found to form a thick fluoride film when anhydrous HF + H<sub>2</sub>O was used under more controlled conditions [31]. Commercial HF-vapor etchers are now available.

**Technics O<sub>2</sub>:** Technics West PE II-A plasma etcher using the recipe O<sub>2</sub> flow rate ≈ 51 sccm, power (P) = 400 W at 30 kHz, pressure (p) ≈ 300 mtorr. Oxygen plasmas are used for stripping and desmearing photoresist, as well as cleaning off other organics. In these tests, the photoresists, Parylene, and polyimide etched at about the same rate, but the graphite was not etched.

**DRIE HF Mech:** Surface Technology Systems (STS) Advanced Silicon Etch (ASE) inductively coupled plasma DRIE tool, using a typical Bosch-process recipe for nearly vertical trenches with high platen frequency [33], [34]: etch for 6 s with ST<sub>0</sub> = 80 sccm, coil P = 600 W at 13.56 MHz, platen P = 12 W at 13.56 MHz, p ≈ 12 mtorr; then passivate for 5 s with C<sub>1</sub>F<sub>8</sub> = 80 sccm, coil P = 600 W at 13.56 MHz, platen P = 0 W, p ≈ 8 mtorr. The etching and passivation steps are alternated. The platen structure has 20 °C coolant flowing through it, while the wafer heats to about 65 °C during an etch. The wafer clamping in this particular etcher, required due to back-side helium pressure for improved heat transfer, is mechanical.

This etcher and recipe are designed for deep, near-vertical etching of silicon. The etch rate of silicon is a strong function of the area of silicon exposed and the trench aspect ratio (depth/width) for deep etches, slowing with increases in each. The etch rate also tends to be noticeably slower at the center of the wafer. In these tests, a higher-than-typical area of ~50% was exposed. Due to a relatively short etch time, only shallow trenches were formed. Experience with several such etchers and chucks (with either mechanical or electrostatic wafer clamping)

shows that the etch rate and profile are highly variable with equipment. Recipes are usually not transferable from one machine to another and can be affected by seemingly small changes in the setup of an etcher.

Photoresist and oxide have been used as masks for DRIE. These tests confirmed very high selectivity of etching silicon over photoresist and silicon dioxide, and also found high selectivity over silicon nitride. The manufacturer recommends not using metal masks. Several metals with volatile fluorides were tested and found to etch slowly with this recipe.

**DRIE HFES:** Surface Technology Systems ASE inductively coupled plasma DRIE tool, using a typical Bosch-process recipe for nearly vertical trenches with high platen frequency: etch for 9 s with  $SF_6 = 130$  sccm,  $O_2 = 13$  sccm, coil  $P = 600$  W at 13.56 MHz, platen  $P = 12$  W at 13.56 MHz; then passivate for 7 s with  $C_4F_8 = 85$  sccm, coil  $P = 600$  W at 13.56 MHz, platen  $P = 0$  W, APC = 60%; with electrostatic wafer clamping.

**DRIE LF Mech:** Surface Technology Systems ASE inductively coupled plasma DRIE tool, using a typical low-platen frequency, stop-on-oxide recipe for nearly vertical trenches [35]: etch for 5 s with  $SF_6 = 125$  sccm, coil  $P = 600$  W at 13.56 MHz, platen  $P = 30$  W at 380 kHz pulsed on and off on the order of 10 milliseconds (the exact timing is proprietary),  $p \approx 28$  mtorr; then passivate for 6.7 s with  $C_4F_8 = 150$  sccm, coil  $P = 600$  W at 13.56 MHz, platen  $P = 0$  W,  $p \approx 28$  mtorr; with mechanical wafer clamping.

**DRIE LF ES:** Surface Technology Systems ASE inductively coupled plasma DRIE tool, using a typical low-platen frequency, stop-on-oxide recipe to greatly reduce notching, for nearly vertical trenches: etch for 8 s with  $SF_6 = 80$  sccm, coil  $P = 600$  W at 13.56 MHz, platen  $P = 30$  W at 380 kHz pulsed on and off on the order of milliseconds (the exact timing is proprietary),  $p \approx 12$  mtorr; then passivate for 5 s with  $C_4F_8 = 80$  sccm, coil  $P = 600$  W at 13.56 MHz, platen  $P = 0$  W,  $p \approx 8$  mtorr; with electrostatic wafer clamping.

**STS 320  $SF_6$ :** Surface Technology Systems 320 parallel-plate RIE tool, using the recipe  $SF_6 = 25$  sccm,  $P = 100$  W at 13.56 MHz,  $p = 20$  mtorr. The driven electrode area is about  $900 \text{ cm}^2$ . This etch targets silicon, silicon nitride, and transition metals with volatile fluorides. Photoresist can be used as an etch mask for this etch.

**STS  $SF_6 + O_2$ :** Surface Technology Systems 320 parallel-plate RIE tool, using the recipe  $SF_6 = 25$  sccm,  $O_2 = 10$  sccm,  $P = 100$  W at 13.56 MHz,  $p = 20$  mtorr. The driven electrode area is about  $900 \text{ cm}^2$ . This etch targets silicon, silicon nitride, and transition metals with volatile fluorides. The addition of oxygen increases the etch rate of the target materials, but also increases the etch rate of the photoresist, which can reduce the selectivity.

**STS 320  $CF_4$ :** Surface Technology Systems 320 parallel-plate RIE tool, using the recipe  $CF_4 = 25$  sccm,  $P = 100$  W at 13.56 MHz,  $p = 60$  mtorr. The driven electrode area is about  $900 \text{ cm}^2$ . This etch targets silicon dioxide, which can be patterned with photoresist or a nonvolatile metal such as chromium. This recipe results in fluorocarbon polymer buildup on the chamber walls and a reduced etch rate over time. A small amount of fluorocarbon residue left on the wafer can affect the later formation of electrical contacts if it is not removed.

TABLE IX  
ETCH RATES OF GOLD DEPOSITED BY THREE METHODS IN TWO  
ETCHANTS (nm/min)

Etchant	Ion-Milled Gold (at 1000 V)	Evaporated Gold (at 0.5 nm/s)	Electroplated Gold (NaAuSO <sub>3</sub> , 40 mA/cm <sup>2</sup> )
Cyanate AU-5	490	680	560
Dilute aqua regia	530	680	1000

**STS  $CF_4 + O_2$ :** Surface Technology Systems 320 parallel-plate RIE tool, using the recipe  $CF_4 = 25$  sccm,  $O_2 = 5$  sccm,  $P = 100$  W at 13.56 MHz,  $p = 60$  mtorr. The driven electrode area is about  $900 \text{ cm}^2$ .  $CF_4$ -based etches typically target silicon dioxide, but this etch was found to etch silicon nitride and silicon even faster than oxide. The addition of oxygen in this recipe increases the oxide etch rate but also increases the photoresist etch rate, significantly reducing the selectivity. The oxygen also reduces fluorocarbon polymer buildup on the chamber walls, yielding a more stable etch rate over time.

**Ion Mill:** Commonwealth Scientific ion mill, ion milling with argon ions at 500 V, current density of  $\sim 1 \text{ mA/cm}^2$ , chamber  $p = 2 \times 10^{-4}$  torr, at an angle normal to the wafer. Ion milling with an inert gas such as argon is a purely physical process, so any material can be etched by this method. The surface of unpatterned S1822 photoresist became wavy after a 10-min etch but retained most of its thickness, so this photoresist may not be a useable masking material.

### C. Etch-Rate Variation

Many reasons for etch-rate variation were enumerated earlier [3], [14]. One of the most significant is the properties of the material, which are a result of the production method and subsequent processing (e.g., annealing). This is apparent by inspection of Table III: materials that are primarily silicon dioxide have widely varying etch rates in 5:1 BHF and other etchants.

As another demonstration of the effect of deposition conditions, gold was deposited using three different methods: ion milling and evaporation, as described in Section II, and by electroplating from a sodium gold sulfite bath at current density of  $40 \text{ mA/cm}^2$ . The etch rates in the AU-5 and dilute aqua regia solutions are given in Table IX. Ion-milled gold, which is the densest, has the lowest etch rate in both etchants. Electroplated gold, known for having a lower density than the bulk metal, has a significantly higher etch rate in dilute aqua regia.

Another example of etch-rate variation with material deposition method is given in Table X. PECVD silicon nitride was deposited using the recipe described in Section II, but with varying flow rates of 2% SiH<sub>4</sub>/Ar. For the lowest silane flow rates, the silicon content is probably lowest, which agrees with the low index of refraction. These films have the highest etch rates in 10:1 BHF. Higher silane flow rates yield increasingly higher indexes of refraction and lower etch rates. The lowest etch rate, 2.5 nm/min, is still much faster than for LPCVD silicon nitride (which has much less hydrogen).

TABLE X  
PECVD SILICON NITRIDE INDEXES OF REFRACTION AND ETCH RATES  
IN 10:1 BHF (nm/min)

2% Si in Ar (sccm)	Index of Refraction	Etch Rate in 10:1 BHF (nm/min)
70	1.83	11
100	1.88	7.3
200	2.03	4.3
500	2.53	2.7
750	2.57	2.8
1000	2.67	2.5

#### IV. ETCH-RATE MEASUREMENT TECHNIQUES

All films were coated onto 100-mm silicon wafers. The transparent films, opaque films, and wafer substrates required different measurement techniques.

Most of the data presented here has not been reported before. In the cases in the tables for which new data was not taken, etch rates were transferred from our earlier paper [3]. In case of discrepancies, the newer data is reported. Most of the ion-mill data for the elements is from Commonwealth Scientific [36] and agrees with our few measurements. This outside data is included due to the usefulness of this tool.

Measured etch rates are rounded to two significant figures.

##### A. Transparent Films

The transparent films in these tests are polycrystalline silicon, silicon dioxide, silicon nitride, alumina, photoresist, parylene, and polyimide. Refractive indexes (RIs) were measured with a Gaertner Scientific ellipsometer at 632.8 nm for most films (the RI of polysilicon is already known). These RIs were used in interferometric thickness measurements using a NanoSpec AFT tool. Polycrystalline silicon was deposited on a wafer that already had 100 nm of thermal oxide to give reflections that could be detected by the NanoSpec.

To reduce the number of samples to be prepared, some wafers were divided into quarters. For wet etches and dry etches with manual wafer loading, the quarter wafers were used. At least three thickness measurements were made before and after etching. Etching was done without masking. For dry etches with automatic wafer loading, a whole wafer was used and five points were measured. The average thickness difference, divided by the time, gives the etch rate.

##### B. Opaque Films and Wafers

The opaque films used in these tests are graphite, the metals, titanium nitride, and the resist pen. Ge and SiGe films, while not opaque, were measured with this group. The wafers are single-crystal silicon, fused quartz, Pyrex 7740, and sapphire. These samples could not be measured by the simpler interferometric method, and a calibrated Dektak II profilometer was used. In most cases, the wafer had photoresist applied and patterned, with about 50% of the wafer area exposed. In a few cases, the resist pen was used to make a mask. For testing the photoresists and resist pen themselves, the photoresist was patterned

and ink was simply drawn onto a wafer known not to etch. For cases in which the etchant is known to rapidly etch photoresist (e.g., in KOH and Piranha), a mask of evaporated Cr/Au was used. For cases in which a silicon wafer is known to etch rapidly (e.g., in silicon isotropic etchant and KOH), the wafers were first coated with LPCVD silicon nitride to protect both sides.

Etches were performed, the photoresist (or Cr/Au) was stripped off, and at least three step heights were measured. The average step height divided by the time gives the etch rate.

For very rapid etches, step heights were measured where the removal of Kapton tape after deposition had left a blank spot on the wafer. The materials were etched until visibly gone and the time recorded. The initial thickness divided by the time gives the etch rate.

##### C. Etch Times

Etch times were selected to give a thickness change typically in the range of 0.1 to 1  $\mu$ m to give good confidence to the results.

As expected, however, many samples etched very slowly. In these cases, the etches were performed for at least 20 min to give a reasonable step height or interferometric thickness change, if any. For the samples measured by profilometer, when it was impossible to see a clear step amidst the roughness, an upper limit of etch rate (e.g., <2 nm/min) is given. This is calculated by dividing the roughness by the etch time.

In some other cases, the etch was so rapid (e.g., photoresist in Piranha and acetone) that it was impossible to tell exactly when the film was gone due to clouds or bubbles blocking view of the film. For these cases, a lower limit of the etch rate is reported (e.g., >92 000 nm/min). This is calculated by dividing the initial thickness by the time at which the clouds or bubbles had dissipated enough to view the cleared wafer.

#### V. ETCH-RATE RESULTS AND DISCUSSION

Etch rates were reported in Tables II–VII. Selectivities can readily be calculated from these data.

Not all materials were etched in all etchants due to time limitations and contamination concerns with the plasma etchers. Certain materials commonly used in micromachining were considered key materials (undoped polysilicon, thermal silicon dioxide, stoichiometric LPCVD silicon nitride, Al/2%Si, chromium, and Si182 photoresist) and were etched in most etchants in order to determine their compatibility. Similarly, certain etches (isotropic silicon etchant, potassium hydroxide, 5:1 BHF, CR-7 chromium etchant, Piranha, ion milling, and SF<sub>6</sub>+O<sub>2</sub>, and CF<sub>4</sub>+O<sub>2</sub> plasmas) were considered to be key etchants and most materials were etched in them.

One of the realities of putting materials in etchants intended to etch a different material is that a clear result is not attained. In many cases materials were roughened, but little etching apparently occurred. The degree of roughening probably depends on the microstructure (i.e., grain size and dislocation density) and thus varies with the method of material preparation.

At the bottom of Table I is a summary of the letter notations used in the tables. In cases where etch rates were not measured by the authors, but the etches are known from experience with

similar materials or from the literature to *work*, be relatively *fast* (faster than  $\sim 500$  nm/min), or be *slow* (slower than  $\sim 10$  nm/min) or have zero etch rate, the notations *W*, *F*, or *S*, respectively, are used. If the film was *roughened* or attacked, the notation is *R*. This may include an upper limit of etch rate (e.g.,  $< 1$  nm/min) if an exact value could not be determined. If the film was *thicker* after being exposed to the etch, the notation is *T*. This occurred with the photoresists absorbing some of the liquid and swelling, and in other cases in which an insoluble compound formed. If the film *peeled* during the etch or rinsing, the notation is *P*. Some etches initially had a very slow etch rate during an "incubation time," and are so noted by *I*. Films that *congealed* are noted with *C*. Films that remained after exposure to the etch, but were *soft* have a notation of *soft*.

#### A. Discussion of the Etches

Intended uses of the etches and specific information about the materials were given in Sections II and III. More general observations are given here.

KOH did not appreciably etch most of the transition metals and the silicon nitrides, and had a zero etch rate for Mo, W, Ni, Pd, Pt, and Au. These might be used in micromachining as structural materials, with the underlying silicon substrate being removed in KOH.

The HF solutions, 5:1 BHF and 10:1 HF, did not appreciably etch most of the transition metals, and had a zero etch rate for Nb, Cr, Ta, Pd, Pt, and Au. The exception is Ti, which etches even faster than the silicon dioxides in 10:1 HF. Thus, Cr, 10 Ti/90 W, or TiN should be used rather than Ti as an adhesion layer in the presence of HF. The metals with a slow HF etch rate might be used in surface micromachining as structural materials, with silicon dioxide being the sacrificial layer.

Piranha removed most metals at moderate to fast rates. Nb, Ta, Cr, Mo, Pd, Au, and 10 Ti/90 W were etched only slowly (Au not at all), allowing the use of Piranha for cleaning with these metals present.

Microstrip 2001 and acetone dissolved the resists tested rapidly. Methanol removed some resists and made a mess of others, causing them to peel or congeal. Isopropanol had much less effect, with the slow removal of resist.

The oxygen plasma only etched the organics, but did not measurably etch graphite.

The STS DRJE's etched silicon and germanium much faster than the oxides, nitrides, and photoresists, making the latter useful as masking layers. The transition metals tested, all of which have volatile fluorides, etched only slowly.

The  $\text{SF}_6 + \text{O}_2$  plasma etched silicon the fastest, the silicon nitrides at a moderate rate, and the silicon dioxides slowly. Moderate etch rates were also measured for Mo and 10 Ti/90 W. Other metals, which do not have volatile etch products, were removed slowly due to sputtering.

The  $\text{CF}_4 + \text{O}_2$  plasma etched the silicon nitrides faster than silicon or oxide, but at only a moderate rate. For the metals tested, molybdenum had a moderate etch rate, and other metals again were removed slowly due to sputtering.

Ion milling can remove any material due to its purely physical nature, and is thus thought of as having roughly the same etch rate for different materials. The data, however, show that

there actually is a wide range of etch rates such that some materials can be used as etch masks for others. Graphite is etched very slowly. The metals Ag, Au, Pd, Cu, and Pt are all etched more than 20 times faster.

#### B. Discussion of the Materials

Changes in the deposition method of pure materials, as well as the addition of small amounts of impurities, tend to produce much greater differences in etch rate for wet etchants than for plasma etching. Comparing the etch rates of the various silicon nitrides in the 5:1 BHF and  $\text{SF}_6 + \text{O}_2$  plasma illustrates this point, with the ratio of fastest-to slowest-etching nitride in BHF being 46, but only 1.3 for the plasma etch. Comparing the etch rates of the many silicon dioxide samples in 5:1 BHF and in  $\text{CF}_4 + \text{O}_2$  plasma gives similar results.

PECVD oxides and nitride etch faster in most etchants than their LPCVD counterparts. Annealing both PECVD and LPCVD oxides reduces the etch rate in HF solutions. Annealed LTO has similar etching properties to thermal oxide.

Unlike sapphire, the aluminum oxide films were etched in most of the wet etchants tested.

Samples of patterned Cr on top of Au were produced to study a possible electrochemical effect. This Cr etched slower in CR-7, but faster in CR-14, as compared to evaporated Cr on Si. In this particular test, Cr on Au peeled off in 5:1 BHF, but we have successfully used Cr as an adhesion layer for Au in surface-micromachined devices in which the Cr was exposed.

Silver grew thicker in AU-5 and in Cu  $\text{FeCl}_3$  200 etchant, perhaps due to the formation of insoluble compounds such as AgI.

Metals deposited by evaporation, in which the atoms forming the film arrive with low energy, tend to etch faster than the denser films deposited by sputtering and by ion milling, with high-incident-energy atoms.

TiN became soft in silicon isotropic etchant. Titanium became soft in KOH. Molybdenum became soft in CR-14. The etch rates reported are based on the difference between the initial thickness and the thickness of the remaining material.

Polyimide stands up to wet etchants better than the resists, but has similar etch rates in the plasma etches. Parylene-C did not etch appreciably in the wet etchants.

#### VI. CONCLUSION

The data in the tables show useful etchants for most materials, as well as indicating in which etchants the materials will survive. They can also be used to calculate selectivities to choose masking layers should photoresist not be sufficient.

For surface micromachining, the traditional system of materials and etchant is a structural material of polysilicon, an underlying layer of polysilicon, a sacrificial layer based on silicon dioxide, silicon nitride, or silicon wafer, and an HF-based etchant. These data show that a number of other materials might be substituted as the structural layer: LPCVD silicon nitride, V, Nb, Ta, Cr, Mo, W, Pd, Pt, Cu, Ag, 90 Ti/10 W, 80 Ni/20 Cr, or polyimide. For bulk micromachining of a silicon wafer with KOH to leave a freestanding film, many of the same materials

might be used: silicon nitride, Nb, Ta, Cr, Mo, W, Ni, Pd, Pt, and Au. Other systems might be used, such as a sacrificial layer of Al etched with Aluminum Etchant Type A, and a structural layer of PECVD silicon dioxide, Ti, or Cr. Another possibility is a sacrificial layer of polysilicon or single-crystal silicon etched with  $\text{XeF}_2$ , and a structural layer of  $\text{SiO}_2$ , Au, or Pt. Many more systems can be deduced from the data.

Alternative materials systems for MEMS will be of interest in the future for many reasons: A previously unused structural material will be desired, for example, for its chemical inertness, for its high or low Young's modulus, for its high or low thermal conductivity, for its low deposition temperature, or for its biological compatibility. These data will aid in selecting appropriate etchants, substrates, and coexisting materials.

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